## **PATENT**

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:

AKZO PATENT

VANDEVOORDE, P. et al.

Serial No.: 09/444.968

Filing Date: November 22, 1999

For: COATING COMPOSITION BASED ON A HYDROXY GROUP-CONTAINING FILM FORMING POLYMER, A POLYISOCYANATE COMPOUND AND A DIOL

**Assistant Commissioner of Patents** 

Washington, D.C. 20231

Docket: ACO2587P1USAX RECEIVED

Examiner: Bissett

Group Art Unit: 1773

<u>CERTIFICATE OF FACSIMILE TRANSMISSION</u> hereby certified that the attached: TRANSMITTAL LETTER IN DUPLICATE;

RESPONSE AFTER FINAL AND CERTIFICATE OF FACSIMILE (17 sheets) is being faxed to (703) 872-9311 to the Assistant Commissioner for Patents

February 10, 2003

## RESPONSE

Sir:

In response to the Office Action of August 8, 2002, Applicants provide the following remarks for entry in this application. This response is filed in accordance with 37 CFR 1.111.

Presently, claims 1-17 are pending, though claims 7-11 and 13-17 have been withdrawn from consideration.

As a first matter, Applicants acknowledge and thank the Examiner for allowing claim 6.

Before addressing the specific rejections involved in this case, Applicants would like to clarify some things with regard to the Examiner's assertions in the present office action.

The Examiner appears to erroneously take the position that a claim directed towards a composition comprising components would include a composition comprising the reaction products of said components. More particularly, the Examiner alleges that a composition comprising a diol and a disocvanate would include a composition comprising the polyurethan being the reaction product of said components.

Applicants are truly perplexed with regard the Examiner's statements regarding Ho. Under the standard of anticipation, Ho does not disclose the present invention. Ho must contain within its four corners a sufficient description to enable one to practice the invention of the rejected claims without undue experimentation or inventive skills. Akzo N.V. v. U.S. Intern. Trade Com'n, 1 USPQ2d 1241, 1245 (Fed. Cir 1986). As set forth previously and as described by the Examiner, Ho does not anticipate the present invention.

The interpretation of the claim language in Ho cannot be expanded to include any and all possible materials the words might read on, only the functional equivalents of what is disclosed in the specification. In re Donaldson Co., Inc., 29 USPQ2d 1845 (CAFC 1994). Nothing in Ho discloses the diol and the diisocyanate as the coating composition.

If there is not a reasonable certainty that the claimed subject matter will necessarily result, a rejection based on anticipation must fail. In re Brink, 164 USPQ 247, 249 (CCPA 1970). In the present instance, nothing in Ho discloses the coating composition of the present invention having AS SEPARATE (NON REACTED) components a diol and diisocyanate....and nothing in the Ho examples results in a coating composition having AS SEPARATE (NON REACTED) components, a diol and diisocyanante.

A first important difference between a composition comprising a diol and a polyisocyanate and a composition comprising a polyurethane lies in their respective viscosities. On the one hand, monomeric diols and isocyanates are usually low in viscosity and serve as diluents for a coating composition. On the other hand, polymers can be highly viscous or even be solid, depending on the molecular weight for a given polymer (J. Bentley, G.P.A. Turner, Introduction to Paint Chemistry, 4th Edition, Chapman & Hall, 1998, p. 131, first full paragraph) (copy attached). The viscosity of a coating composition governs its application properties (Bentley et al., p. 93, last paragraph, p. 94, 1st paragraph) (copy attached). In the present case and as set forth above, Applicants respectfully request that it the Examiner maintains that the Ho reference anticipates the present invention, she specifically set forth, within the confines of a coating composition, how Ho could be applied to a substrate in the intermediary stage.

A second difference between a composition comprising a diol and a polyisocyanate and a composition comprising a polyurethane lies in storage stability. A composition comprising a diol and a polyisocyanate increases in viscosity after mixing due to the reaction of the components and the composition thus has a pot life (Table 3 on p. 17 of the application). Conversely, a polyurethane is depleted of isocyanate groups and is storage stable for a long period of time without any change in properties. A skilled person is aware of the importance pot life has as a property of two pack paints (Bentley et al., p. 100, paragraph titled "Drying by chemical reaction between ingredients in the paint").

A third difference is that a diol is a compound of a specified formula characterized by a special arrangement and connectivity of atoms, and by the presence of functional groups, namely two hydroxyl groups. In the reaction with polyisocyanate the hydroxyl groups are converted to urethane groups. In the light of the foregoing explanation it should be clear that the urethane resin thus prepared no longer comprises the diol as a compound with hydroxyl groups, but a polymer having urethane groups.

Thus, a skilled person is very well aware of the differences between a composition comprising a diol and a polyisocyanate and a composition comprising a polyurethane. For coating compositions in particular, these differences are of great importance. Accordingly, a skilled person will not interpret a claim reading on a coating composition comprising a diol and a polyisocyanate as including a coating composition comprising the polyurethane being the reaction product of said diol and said polyisocyanate.

Specifically, claims 1, 2, 4, 5, and 12 stand rejected for alleged anticipation by Ho. Applicants respectfully maintain their traverse of this rejection for at least the reasons set forth in previous responses, the remarks made above and for the following reasons.

Ho relates to water-based reactive two-part polyurethane compositions comprising a part A and a part B (col. 2, II. 18 – 22), and to non-aqu ous based compositions comprising a part A and a part B (col. 4, I. 40 – col. 5, I. 34).

For aqueous-based two-part compositions, part A is an aqueous composition and optionally comprises

- (a) urethane prepolymer having hydroxyl functionality and, optionally,
- (b) a water-reducible first acrylic polymer having at least one hydroxyl group, and optionally further components (c) (h) (col. 2, l. 38 col. 3, l. 41).

Part B comprises a crosslinker selected from polyisocyanates, blocked polyisocyanates, and mixtures thereof (col. 3, II. 36 – 41).

The Examiner erroneously alleges that in Example 8 and in Table 5, col. 38 Ho discloses <u>a coating composition</u> to be used as clearcoat comprising 2-butyl-2-ethyl-1,3-propanediol, an isocyanate, and two polyol compounds.

Conversely, Example 8 actually describes the preparation of a PUR (polyurethane) dispersion used as **component** (a) in part A of the two-part compositions cited above (col. 30, II. 29 – 40). In Example 8 reference is made to the manner of Example 1. In Example 1 it is described that the reaction was allowed to run to completion [no residual isocyanate was detected by FTIR] (col. 29, II. 14 – 16).

Ho discloses that the compositions formed by combining part A and part B can be used as coating composition (col. 23, II. 48 – 50). However, Ho does not disclose that the components used to prepare a polyurethane dispersion, such as the components of Table 5, can be used as a coating composition. Applicants again respectfully request that if the Examiner maintains her objection, that she specifically demonstrate how, under the established rules for defining a coating composition, how the components in Ho (not the resultant polyurethane dispersion) can be used as a coating composition.

More particularly, we further maintain that the components of Example 8 in Table 5 cannot be used as coating composition as such, due to inherent instability, since they exotherm to 85°C. On the other hand, if one considers the polyurethane formed after the reaction was allowed to run to completion, it is

d plet d of isocyanate (col. 29, ll. 14 - 16) and would not have this problem. Applicants request scientific support for the Examiner's contention that inherently unstable components can be used as a coating composition.

Ho does not disclose a coating composition according to instant claim 1. Accordingly, claim 1 and all depending claims of the current application are novel over Ho.

Assuming arguendo that Ho disclosed even the polyols of instant claim1, the Examiner quotes the hydroxyl equivalent weights of the polyols used by Ho for the preparation of a polyurethane dispersion in Example 8. We presume that the Examiner is trying to demonstrate that the polyols used by Ho in Example 8 in the preparation of a PUR dispersion meet the film-forming polymer of claim 1 of the instant application. We would like to respectfully remind the Examiner that a hydroxyl value expressed in mg KOH/g is different from a hydroxyl equivalent weight (Nylen, Sunderland, Modern Surface Coatings, Interscience Publishers, 1965, p. 58, copy attached).

Polyol IV of Example 8 of Ho has a hydroxyl equivalent weight of 98 g/equivalent. 1 g of polyol IV thus contains 1/98, i.e. 0.0102 equivalents of hydroxyl groups. The equivalent weight of KOH is 58,100 mg/equivalent. In order to calculate the hydroxyl number of polyol IV one has to multiply the number of hydroxyl equivalents in 1 g of polyol, i.e. 0.0102, by the equivalent weight of KOH, i.e. 56,100. This corresponds to a hydroxyl number of 572.4 mg KOH/g of polyol IV of Example 8 of Ho.

Polyol I has a hydroxyl equivalent weight of 72.9 g/equivalent. This corresponds to a hydroxyl number of 769.5 mg KOH/g.

Thus, neither polyol IV nor polyol I, used by Ho meet the film-forming polymer of instant claim 1.

The Examiner also rejects Claims 1 – 3, 5, and 12 for alleged anticipation by Mayer. Applicants respectfully traverse this objection.

Mayer discloses a coating composition comprising a component (I) containing a binder and a component (II) containing a polyisocyanate. Component (I) comprises an acrylate copolymer and optionally a polyurethane

Feb 10 2003 16:58

resin (col. 15, II, 56 - 67). The polyurethane resin may be prepared by r acting isocyanate-containing prepolym rs with compounds, which are reactive toward isocyanate groups (col. 12, II. 13 - 16). 2-butyl-2-ethylpropane-1,3-diol is mentioned as a suitable compound which is reactive with isocyanate groups (col. 13, II. 15 – 23).

Mayer does not disclose a coating composition comprising a diol according to the formula of current claim 1.

Therefore, and in view of the explanation above, current claim 1 and all depending claims are not anticipated by Mayer.

Finally, the Examiner rejects claim 3 as obvious under 35 USC § 103. Applicants respectfully traverse this objection.

Claim 3 stands rejected for alleged obviousness over Ho. Ho notes the possible addition of a polyacrylate polyol to part A of the two-part composition: (col. 16, II. 18 - 34). Part A of the two-part compositions of Ho can comprise components (a) - (h) for water borne compositions (col. 2, l. 38 - col. 3, l. 18) and components (a) - (g) for non-aqueous compositions (col. 4, I. 39 - col. 5, I. 14). The optionally present polyacrylate polyol corresponds to component (b) of part A of said two-part compositions.

However, none of the components (a) – (h) and/or (a) – (g) of part A corresponds to the diol according to the formula of instant claim 1.

Furthermore, as set forth above (concerning the difference between a composition comprising a diol and a polyisocyanate and a composition comprising a polyurethane which is the reaction product of the components), it should be realized that part A of Ho's two-part composition does not comprise a diol according to the formula of instant claim 1, even if a polyurethane corresponding to component (a) is the reaction product of such a diol and Isocyanates.

Accordingly, including a polyacrylate polyol in part A of the two-part composition of Ho does not lead to the composition of instant claim 3. Therefore, claim 3 is not obvious in view of Ho.

AKZO PATENT

In view of the amendments and remarks herein, the papers submitted previously, the pres nt application is believed to be in condition for allowance, which action is respectfully requested.

Respectfully submitted,

Feb 10 2003 16:58

Joan M. McGillycuddy Attorney for Applicants

Reg. No.: 35,608

Akzo Nobel inc. Intellectual Property Dept. 7 Livingstone Avenue Dobbs Ferry, New York 10522-3408 (914) 674-5463

> **FAX RECEIVED** FEB 1 1 2003 GROUP 1700

Introduction to

Paint Chemistry

and principles of paint technology

I. Bentley

and

itath farmerly with ICI Paints Skaugh, Berlashira, UK G. P. A. Turner

Fourth edition

Landon - Weinheim - New Yark - Takya - Meibenana - Madass CHAPMAN & HALL

JOIN US ON THE INTERNET VIA WWW, GOPHER, FTP OR EMAIL:

WWWV: http://www.thamson.com GOPHER: gcpher.thamson.com FTP: ftp.thomson.com EMAIL: findil@klask.thamson.com

A Service of ICPP

3

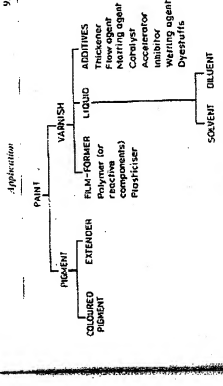


Fig. 7.1 Compresents of paints.

Pigment. Any fine solid particks that do not dissolve in the varnish. If they do not provide colour they are called extender particles. Extenders are much cheaper than prime pigments and can earry out many useful funclious, e.g. improvement of adhesion, ease of sanding and film strength.

film-former. When the conting is dry the film-former is a polymer, but in the wer sample it may be the chemical ingredients, only partly polymeric, which react to form the final dry polymer.

Liquid. Some of the liquids of the paint are often withhold from the paint container and supplied separately as a thinner. Users add thinner to adjust

Additives. Small quantities of substances utiled to earry out special jobs, such as the improvement of surface appearance. the consistency to their requirements.

# Application

the excess of which is then utlawed to drain off, or the correct quantity of These are the basic ingredients, but much care in formulation will be required to practuce a paint that will be easy to apply under changing conditions and pleasant to look at when dry. It is not within the nims of this book to discuss the methods of application that are available. They are well described elsewhere (see Appendix A). Suffice it to say that the paint may be put on by breshing, by roller, by a whole variety of methods of spraying (compressed air, airless, chetrostatic and aerosol spraying), by dipping, chetrodeposition, curtain-exating and flow-exating. Either the article is immersed in the paint, paint is applied to the article and must not drain, 'run' or 'sag'.

In all cases, any irregularities in the wei film cansud by the multhod of application must flow out to knive a smooth surface.

- used to describe all clear avoidmishes. Enamels are normally thermosetting charcoat is used for unpigmented toatings applied over metallic paints. The faish or topoon, meaning that it is the fast coat to be applied and the one planic solution paints or varnishen, dut the tertu is sometimes (confusingly) If the pigment is omitted, the material is usually called a varulsh. The term numented varnish . the paint .. is sometimes called an enamel, languer, seen when the contod object is examined. Lasquers are normally thermopaints, hard, with a superficial resemblance to vitreaus enamels.
- Paint applied before the topcoat is culled an undercuat. Some undercoats and related finishes may be briefly defined as follows:
- . Filters or stoppers are materials of high solid content, used to fill holes and deeper irregularities and to provide a level surface for the next coat.
- to prevent absorption of later coats by pornus surfaces and to give corrosion resistance over metals. As their name implies, they prepare the Primers are applied to the filled or unfilled surface, to promote adhesion, base material for further paint application. Special pigments improve the anti-currosive properties.
- body of the paint film. level out minor irregularities in the substrate and They are easily rubbed unrooth with abrasive paper. They provide the Surfacers, or underenats in decorative house juinting, are highly pigmented materials containing large quantities of extender (see below). must stick well to primer and topooat.
- Primer-surfacers are surfacers that can be applied direct to the object's surises (the substrate), and fulfit both of the functions above in one coat. Scalers are clear or pigmented materials applied in thin couts to prevent the passage of substances from one coat of paint to another or from the substrate into later coats. They can be required to improve adhesion
- All these materials are formulated on the principles described above. These are illustrated in Fig. 7.1, which also lists some of the minor ingredients of a pain:. Some of the terms in the diagram used a little further explanation at this stage:

heiween civils, where this is otherwise weak.

pigment and titnsformer in a diluent. Thus the third basic ingredient of pann is a liquid. Often the film-former liquid mixture is called the refliche

or the pagment.

ous, the film-former covers and protects the startage, decorating it by giving is gloss or theen. It is difficult to attach contings that are not fluid to any but the simplest of surfaces: those that are flat or gently curving. The fluidity of paint permits penetration into the most intricate creviers. It is achieved by dissolving the film-former in a solvent, or by collected suspension of bath

ilm-former of binder, which will be a resin or polymer, to hind together the pigment particles and hold them on the surface. If the pigment is left protects, but it tubs off. So most paints contain the second ingredient, the

Paper first principles

2

Pubir first principles

5

hese reasons, the paint formulator chaoses the ingredients to give the highest

take the polymer out of solution (if this is practicable and suitable) and is necessary to the polymer molecular weight down since, as will be wen in Chapter 9, this reduces the viscosity. Alternatively (Chapter 9 again), it make a dispersion or 'emulsion' paint.

# Dry film properties

cation presents problems which influence the formula. After application comes drying, but before this is considered, the properties required from the dry film An cuttine paint formula has been described and it has been shown that applimust be discussed, since these influence the chrise of method of drying.

beneath, be hard enough and sexistic enough for the purpose for which it is used and must retain most of its protective and desociative properties for a long period. The paint should be capable of repair or renovation. Let us If the dry paint film is to be a useful one, it must stick to the surface now take these requirements separately and see how they are achieved.

Adhesion

another. The strength of this attraction varies greatly with the atoms We have seen that most molecules or atoms have some attraction for one concerned, but all intermokenlar attractions have one thing in common: they operate over computatively short distunces (<10 mm) and become wester the farther upuri the atoms are within that short range. At distances above I am their contribution to adhesion is negligible.

), which limit considerably the choice of solvents for paints to be face tension  $(
u_e)$  of a smooth solid surface is a numerical measure of the cuse of difficulty of werting it. It is equal to the linguest surface transion possessed by any liquid that will sprend spontaneously when placed on it. If a paint is to wet the surface, it must have a surface tension equal to or lower than the critical surface tension of the solid. Physics can have very low critical surface tensions (e.g. polyteirafluoroctliene, PTFE = 18.5 nM m 1; polytelliene = Since these attractive forces are the ones that make things stick together, the first problem is to get the molecules within range, i.e. the paint film inust 'wet' the surface, displacing air and all the other adsorbed materials. The critical surapplied to them (multads of overcaming this are disclosed in Chapter 17). 31 alvin

but even the thinnest possible tayer of adsorbed oil or grease can dominate leading to wetting problems. Even if the paint wets the concominated surface. Luxsely adhering grease, dirt or rust is a menace since, Chean metals generally have higher critical surface tensions (>73 nM m the surface (e.g. & for 'clean' implies cans has been 38 nMm 1).

To keep the paust manageable at high solids, the formulator has to keep solids' possible. the first prablem is to make the paint easy to handle while it is being applied. Different application methods require paints of different consistency. but in all cases the principle is the same: the greater the content of dissolved

So far so good. The difficulty bes in the next stage. Must methods of correct annum of paint, which must flow at first to remove the irregularities, spruy motile, roller supple and so on. These are the methods that apply the and then stop flowing, to prevent 'rumning' or 'sagging' of paint on vartical surthe balance of the types of liquid used, the formulator brings the consistency of application leave some irregularities in the wet film surface: brush marks, polymer in a paint, the more viscous it will be. The weight percentage of involatile material found in the paint is known as the solids content or solids of the point. Fifther by adjusting the solids of the paint, or by altering viscosity of the paint to that required.

faces. This change in the rate of flow is usually brought about by exaporation Flow dwindles until it is scaredly occurring at all. Then the drying mechanism takes over to set the film. A very careful balance of solvents is required to do of salvent, causing a use in 'solids' and bence a thickening of the consistency. this satisfactorily and more will be said about this in Chapter 9.

characteristics, so that it is fluid while being agriated, but thickens up twee a An alternative method, often used in conjunction with solvent evaporation, is to archade some material in the pain which gives it abnormal viscosity period when the unitation stops. This type of paint is said to he thixotropic.

must drain evenly and not so rapidly that the film thickness becomes too If the article is allowed to drain after being coated, as in dipping, then it tow. If possible, thicknesses at the top and bottom of the article must vary only slightly, in spite of the downward drain of paint. Again the same More detail is given in Chapter 10.

principles must be used to slacken the flow-

which evaporation excurs - that is large compared with the droplet's tion as that leaving the speay gun. The paint is broken up into thousands of fine droplets as it leaves the gun, each droplet presenting a surface - at volume. A great deal of liquid can be lost and this must be taken into account With spray application in particular, the position is further complicated by the fact that the paint reaching the surflice closs not have the same composi-

beam, the world stocks of which are fimited and must be conserved. For all with higher 'solids'. Increasingly thinners other than water are all seen as hazardous to sume degree and undesiruble; also legislation is being enauted to reduce the amount of organic material which can be exhausted into the ennscapence of the princess. They are constantly liviking for new materials which will contain less wasted material; in other words they wunt paints amosphere, thereby polluting it. Furthermore, solvents come from petro-It is necessary to point out that this loss of solvent all of which has been paid for ... is only accepted by the paint users on sufferauce, as an inevitable in formulating the paint.

Paint: first principles

The principle of keeping the paint stable by havingone reactant outside the can is quite general. Chemicals not normally in the air can be introduced into air in a confined drying chamber. This technique is used in the 'vapour curing' of isocyanate paints (Chapter 15).

Alternatively, the reactive ingredient can be in the substrate before it is

coatted, or in a previous coating on the substrate. This principle is used in the connect process for cuting unsaturated polyester coatings (Chapter 16). The application of these separation techniques is kinited only by the chemistry of individual paint systems and the imagination of paint chemists.

# Drying by chemical reaction between ingredients in the paint

must all be in the paint. This paradox is resolved either by separating the reactive ingredients in two or more containers and mixing just before use by choosing ingredients which only react at higher temperatures or what is known as a two-pack paint. Two-pack paints are less popular than their ready-mixed equivalents, because measuring is required before period after mixing during which the stowly in the can, but much more quickly on a surface once the solvent has is used, the chemical reactants may be sticky, low nolecular weight polymers, or they may be simple chemicals. The reaction ants must mu react until the paint has been applied, yet (in this method) they when exposed to radiation of some form. The fermer method produces remains usable (the 'put life'). Sometimes two packs are avoided by the reactants with solvent that reaction proceeds only very Obviously the paint must remain chemically stable in storage, and the reacgone. Here the paint is not really slabbe, but a tolerable 'shelf life' is obtained mixing and because of the limited ornchices a cross-linked polymer Whichever method diluting Paint

pinouses a cross-time, projing.
Industrial stoving enamels and auti-corrosive (wo-pack epoxy coatings dry
by chemical reaction of their ingredients.

A summary of the properties associated with the drying machanisms is given in Table 7.1.

# Relative merits

It has been shown that paints are simple in outline, but complex in operation and formulation. How can the best choice of the many ingredients available be match? The choice of pigments and solvents is discussed in the next two chapters. This summary is concerned with the film-former.

Cross-linked or not cross-linked? There are points in lawour of both alternatives. The tracket, trops durable and solvent-resistant films contain cross-linked pulymers, and these are applied at higher solids. If they are to be matched in all respects except solvent resistance, linear polymers of high molecular waight must be used. For solution polymers,

**ADÚNICIA POR A** and polyener entidired polyurchane None mode Two-pack or Hefinish 2K. seid מאכנטניטן אותנותו bros ristalboow games अध्यक्ष stro:page: convicus (specialion בתעונה ואוכם ACIDATED S ecij jan sard cas page %001-0¢ moneuper Desired. 141) j boskuil EBRIODIS SVIIGURDINE ASSOL S Medium to tugh Chemical featies CRIP DO SOULLY DAN INSTRUCTOR Staving and 901104 statistics at hobbe ententhoow REEL TUR (c) chemicals HEL ALIEN Customated bothanta וינסוסטו נוסח 201 THE OT DODON bolithe lipw (d) chemicals Cans must be One-pack polyurchanus 2.50 No.1 MOT ಶಿರ್ಮಗಳಗಾಗಿ ACLUSTED. begut and (s) an, ppgqq %(0111-15 SIMULTING AUTOR AND STREET Soling light SECTION DIGO menopour (,M%Garting souly artinatoraci med meg Agist of multivity HEREITAN PERMENSI Chine thust be ACEN PERMIT -MCRS pissinop (ums;nus) DING HONOKERILO (Izon) mon applicable %01~0v अवस्थाः श्वासाः COTEG JOINA រដ្ឋម៉ូកែ-ពាបប៉ែងស្រី 122017 incultant entitional Ciond (protes: ~2.C. 1247 10% (monaros) missiones Lingus पक्षिम ratio ban wotalicaonti/ poog 2,00 ISE -Cood 4655-131 ,WOL (CE) WUGGE LEDOW. Summed पक्षक ai manol amiriadina Duggspad क्षां स्टब्स् mit to arying. (1899 ถน) and broader endurer4 umumin. रिक्षांट व्यू दृःरे to seed type of \*PHCS Mold W Method Handbag

Table 7.1 Effect of drying process on properties of paints

day with

temperature (usually 25 °C). However, viscosities read directly in viscosity units are obtained easily if the viscometer contains paddle or disc, mechanically driven to make the liquid in an enclosed space. The force tending to twist the shiftnessy part of the apparatus in contact with the liquid is usually measured and the scale is calibrated in poises or Pas. Comparisons between measurements made in different types of instrument can then be made. Another advantage is that either the rate of shear fproportional to the speed of rotation) or the stress (proportional to the driving force) can be varied, depending on the instrument.

The second category of instrument may alternatively involve a ball falling through a liquid, or rolling in a tube containing the liquid. The time taken for movement over a given distance is measured, but conversion of the answer to poises is easily done. The liquids need to be at least partly transparent to use this type, of measurement. Another type, more suitable for very viscous liquids, involves the rise of a hubble in a tube containing the liquid. The time of rise in seconds can be converted to viscosity units. This is typically used for usin solutions and frequently in restin processing. For viscous liquids there is also a method in which the vibration of a real is damped by the liquid, the effect being related to the viscosity. Results can be obtained in poises or Pa s.

# Farts and theory

There are three essential facts concerning the viscosity of a polymer solution. Let us consider their in turn.

these forces may be weak compared with those operating in, say, a sodium chloride solution, they operate over a veny hing length of molecule those forces to come into play. In addition, there is the possibility of Early in this chapter it was stated that polymers do not have a solubility limit become they cannot crystallize. Nevertheless there are forces of altraction operating between the pulymer molecules in solution. Though and there are frequent encounters with other long molecules, enabling simple mechanical tangling, as with pieces of string. Since both factors have been quoted in Chapter I as causes of increasing viscosity, it is not surprising that even comparatively low concentrations of polymer can cause emisiderable thickening of simple liquids. As the 'safak' of the solution increase, the enciveners and entanglements between molecules become more frequent and as the viscatity increaves, e.g. solution viscosities for RS | second nitrocellulose (Chapter 11): 12%, 0.1 Pas; 20%, 1 Pas; 10%, 18 Pas. Eventually the solution becomes so viscous that it cannot So, although there is no limit at which the solution becomes saturated, be used for paints unless further layind is added. At higher solids' still, the solution alnust creases to flow and might be mistaken for a solid

Sofvent properties

there is a limit at which the adultion becomes too viscous to use. No precise figure can be quoted for this, since it depends upon the use.

Since we have the same total weight of polymer in both solutions, there acrylate, in the same solvent at the same level of soluls, the more viscous reinforced by chemical attraction outweighs the reduced number of the liquid. However, to double the molecular weight and halve the if we have two solutions of the same pulymer, e.g. polymethyl methsolution will contain polymer molecules of higher molecular weight. are fewer polymer molecules present in the high molecular weight polymer solution, but they are longer. The increased opportunity for entangleanent molecules. All the attractions between pulymer molecules in a useful solution are continuously forming and breaking with the movement of the molecules. If they were permanent, the solution would not flow, since the polymer molecules would form a semi-rigid, reinforcing network in number of undecutes involves making a number of these temporary associations into permanent chemical bonds. The network is not made rigid, but it is a good deal less flexible and less amenable to being deformed in flow than beloto. Thus high muleculur weight polymers give more vixens volutions, c.g. 10% solutions of PVA of malecular weight 15000, 2.5 mPn s; 73000, 12mPas; 160000, 58mPas.

If we take a given sumpto of polymer of fixed molecular weight and dissalve it at the same 'solids' in a variety of 'true solvents', the viscosities of the solutions will be proportional to the viscosities of the original solvent, e.g. 12% polystyrene in methyl eithyl ketone (0.4 mPas), 40 mPas, in eithyl benzene (0.7 mPas), 160 mPas, in o-dichlorobearene (1.3 mPas), 330 mPas. This is important, because we can reduce the paint viscosity without lowering the solids or polymer molecular weight, simply by changing to a tess viscons solvent, if a xuitable one is available.

Solvents in the middle 80% of the polymer's parameter range are almost certainly good'solvents. Good solvents dissolve the polymer at all concentrations. Some liquids (paor'solvents) give solutions at certain concentrations, but further dilution precipintes the polymer. In these quasi-volutions, polymer molecules collect in clusters, rasing the effective molecular weight of the polymer and giving an abnormally high viscosity. It is worth noting here that if good solvents in a salution are partly replaced by non-solvents, or if poor solvent is used instead, the viscosity can rise. This huppens when the average solubility parameter of the mixture moves to the extreme of the range for the polymer and clustering of polymer molecules occurs. This is the stage prior to precipitation.

Finally, a word about the viscosity of emulsions. If a polymer is fully emulsified in a liquid and no part of it is in solution, then the molecular enaughements and associations do not occur. Consequently the emulsion viscosity intat he independent of the molecular wright of the dispersed

# MODERN SURFACE COATINGS

a textbook of the chemistry and technology of paints, varnishes, and lacquers

5

# PAUL NYLÉN

Professor exter. in the Royal Institute of Technology, Stockholm; Formerly Head of the Central Research Laboratory of the Swedish Palas and Varnish, Industry

and

1965

# EDWARD SUNDERLAND

Research Chemist, AB Alfort & Cronhalm, Stackholm

INTERSCIENCE PUBLISHERS a division of John Wiley & Sons Ltd. London . New York . Sydney

### Published by Chapman & Hall, 2-6 Boundary Now. London SE1 8HN, UK

Chapman & Hall, 2-6 Boundary Row, London SEI 5HN, UK

Chapman & Hall GmbH, Pappelaltee 3, 59469 Weinheim, Germany

Chapman & Hall USA, 113 Fifth Assense, New York, NY 10003, USA

Chapman & Hall Japan, ITP-Japan, Kyowa Baikling, 3F, 2-2-1 Hirakawacho, Chiyoda-ku, Tokyo 102, Japan

Chapman & Hall Australia, 102 Dodds Street, South Melbourne, Victoria 3205, Australia

Chapman & Hall India, R. Scinadri, 32 Second Mam Road, CIT East, Madras 600 035, India

First edition 1987 Second edition 1980 Trard edition 1988 Reprinted 1990, 1991, 1993, 1995 Fourth edition 1998

4; 1967, 1980, 1988 G. P. A. Turper; 1998 J. Bentley and G. P. A. Turper

Typeset in 10/12pt Times by Academic & Technical Typesetting, Bristol, UK Printed in Great Britain by St Edmundsbury Press, Bury St Edmunds, Suffelk

ISBN 0-412-72320-4 (HB) - 0-412-72330-1 (PB)

Apart from any fair dealing for the purposes of research or private study, or criticism or review, as permitted under the UK Copyright Designs and Patents Act. 1988, this publication may not be repredited, stored, or trunsmitted, in any form or by any means, without the prior permission in writing of the publishers, or in the case of reprographic reproduction only in accordance with the terms of the beences issued by the Copyright Licensing Agency in the UK, or in accordance with the terms of the location outside the UK. Enquiries concerning Reproduction outside the term stated here should be sent to the publishers at the London address printed on this page.

The publisher makes no representation, express or implied, with regard to the accuracy of the information contained in this book and cannot accept any legal responsibility or liability for any errors or omissions that may be made.

A catalogue record for this book is available from the firtish Library

⑤Printed on scid-free text paper, manufactured in assortance with ANSI/ NISO Z39.48-1992 and ANSI/NISO Z39.48-1984 (Permanence of Paper).

48

### DRYING OILS

saponification is complete, the excess of alkali is titrated with standard aqueous hydrochloric acid in the presence of an acid/base indicator. Concurrently with this determination a blank test is carried out under identical circumstances, but in the absence of the sample. For details see B.S. 684-1958 and ASTM D 555-61.

As a measure of the content of esters in a substance, the ester value is sometimes quoted, this being the saponification value minus the acid value. In the presence of labile compounds such as lactones, however, the ester value becomes meaningless as a measure of ester content.

Unsaponifiable matter is oil-soluble matter that, under the test conditions, is not converted to insoluble scaps by potassium or sodium hydroxide but is soluble in the fat solvent, e.g. petroleum ether or diethyl ether used for extraction. Included in unsaponifiable matter are, inter alia, hydrocarbons, higher aliphatic alcohols and sterols.

Hydroxyl value is a numerical measure of the content of free hydroxyl groups in an organic substance. It is expressed as the number of milligrams of potassium hydroxide equivalent to the hydroxyl content of 1 g of the substance.

In the standard method the sample is reacted with a pyridine solution of acetic anhydride which reacts according to the equation

After acetylation is complete the excess anhydride is hydrolysed to acetic acid which is determined by titration with standard alcoholic potash solution.

Iodine value is a numerical measure of the content of ethylenically unsaturated compounds in an organic substance. It is expressed as the quantity of halogen, calculated as weight units of iodine, which 100 weight units of the substance take up under defined conditions, in other words, as the weight per cent of iodine.

A weighed quantity of the substance is dissolved in an inert solvent such as chloroform or carbon tetrachloride, and a known quantity of a reagent containing some form of free halogen (Table 3.12-2) is added in excess. After a definite reaction time a solution of potassium iodide is added and the iodine liberated by the excess of halogen is titrated with standard thiosulphate solution, using starch as indicator. A blank determination is carried out concurrently under identical conditions but omitting the sample. For details see B.S. 684-1958, ASTM D 555-61, and Gardner and

Sward.7

There are a number of methods for the determination of iodine value which differ in the type and quantity of halogon used, in the reaction time,